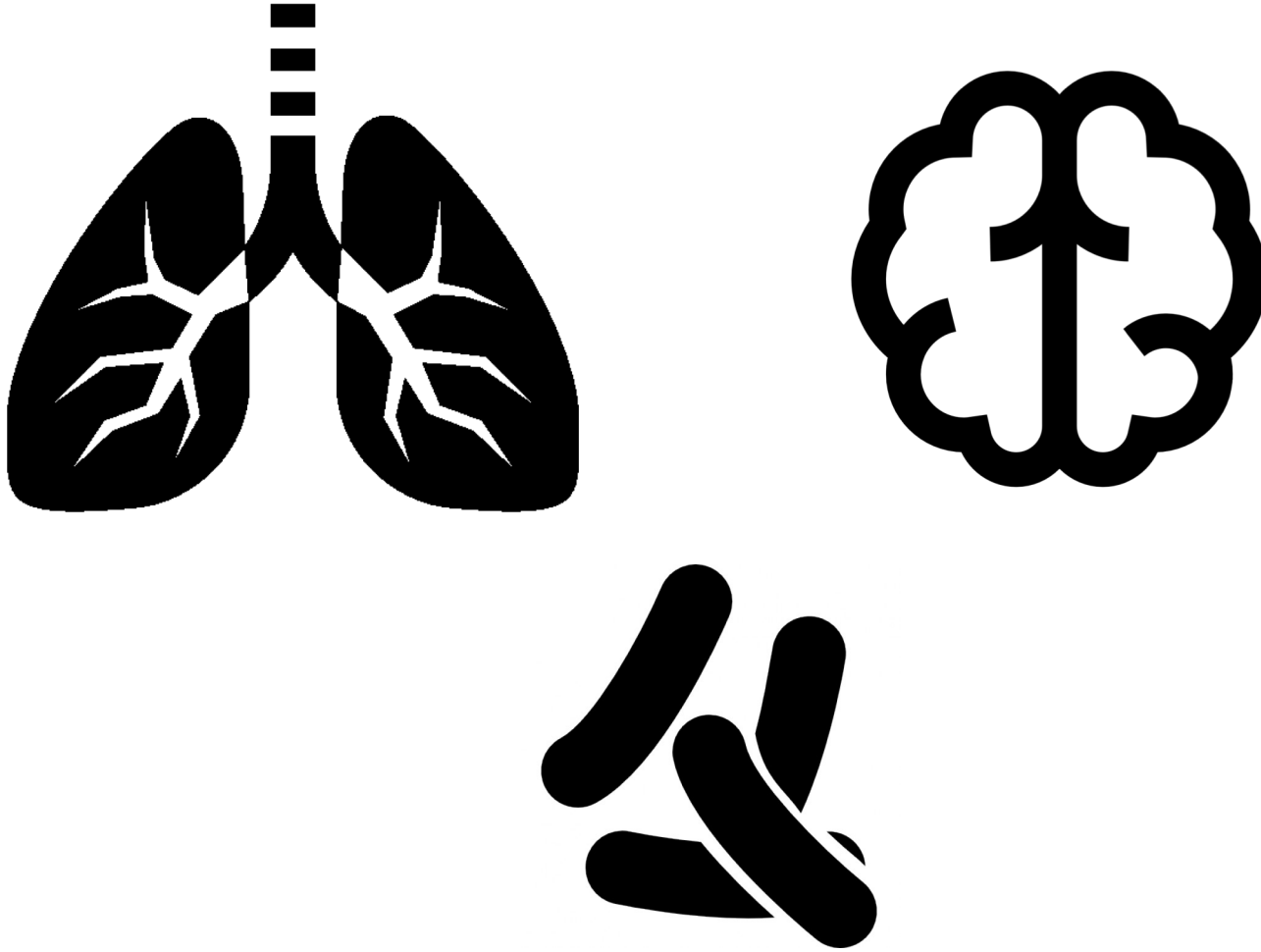


Take A Deep Breath: Lung Infection and Cognition in Down Syndrome



Disclosure

Our lab has received funding from the following:

- **Jerome LeJeune Foundation**
- **Jayden DeLuca Foundation**
- **Celgene, Inc.**
- **American Heart Association**
- **Linda Crnic Institute for Down Syndrome**

Disclosure



Morbidity and Mortality in Persons with Down Syndrome

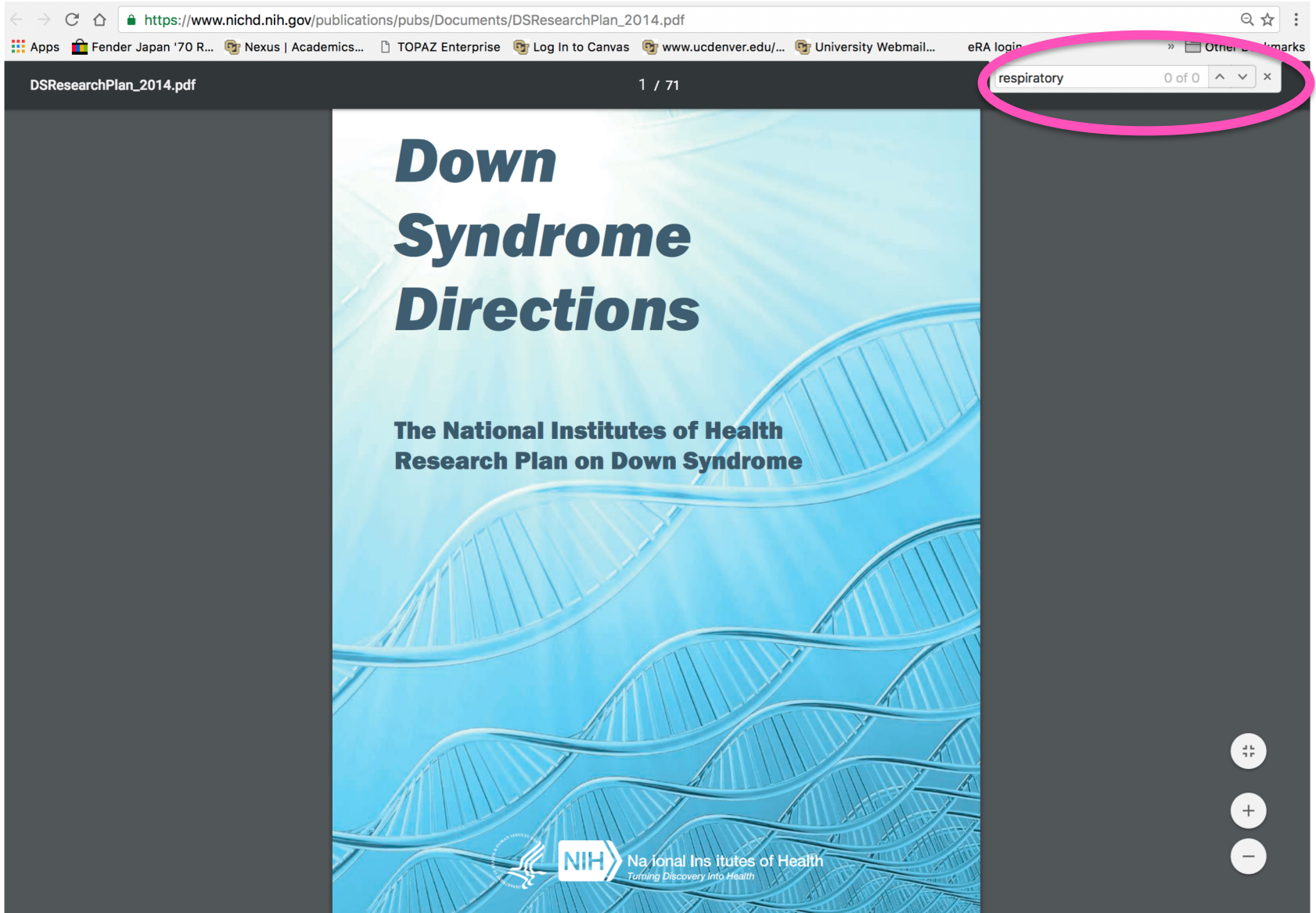
Manifests Predominantly as Infectious Lung Disease

TABLE II. Main and Contributing Causes of Death (%) Over Three Time Periods (1969–1979, 1980–1990, and 1991–2003) in Individuals With Down Syndrome, Divided into Two Age Groups (<1 year, 1-year) and Levels of Significance for Changes Between the Periods 1969–1979 and 1991–2003 (Percentages Will not Add to 100% Since Each Individual Can Have Multiple Diagnoses)

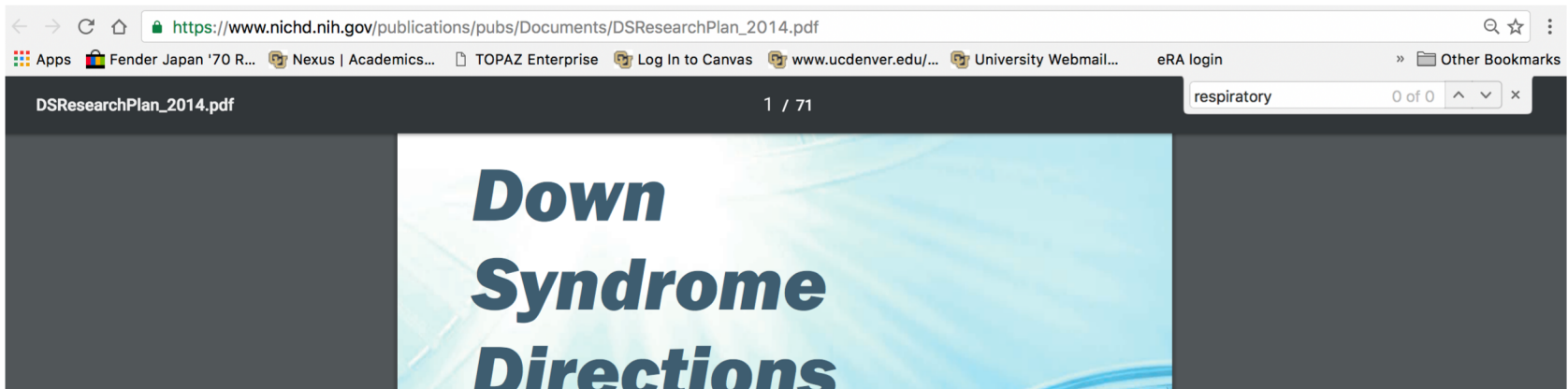
Age	P1		P2		P3		1969–2003	P1 vs. P3	
	1969–1979		1980–1990		1991–2003			P<	P<
	<1 year	1 year	<1 year	1 year	<1 year	1 year		<1 year	1 year
Infectious diseases (incl. pneumonia)	38.3	54.3	27.6	52.0	30.0	57.1	51.2	NS	NS
Pneumonia	31.0	44.0	7.6	41.7	7.1	48.7	40.9	0.001	NS
Congenital heart malformations	74.3	34.6	75.2	26.1	51.4	11.3	29.4	0.001	0.001
Circulatory disease	2.3	17.3	16.2	20.6	35.7	33.3	24	0.001	0.001
Dementia	0.0	0.3	0.0	6.6	0.0	23.3	11.2		0.001
Epilepsy/seizures	0.0	5.9	1.0	9.1	1.4	15.3	9.6	NS	0.001
Atherosclerosis/ischemic heart disease	0.0	6.8	0.0	7.6	0.0	9.7	6.9		NS
Central nervous system disease	0.0	3.2	1.0	7.1	1.4	7.6	5.4	NS	0.004
Gastrointestinal disease	3.4	6.8	3.8	5.6	5.7	4.8	5.2	NS	NS
Malignancies, all	1.7	5.7	1.9	5.3	0.0	2.9	3.7	NS	0.032
Leukemia	1.7	4.3	1.9	3.3	0.0	1.1	2.2	NS	0.001
Solid tumors	0.0	1.4	0.0	2.0	0.0	1.8	1.5		NS
Other congenital malformations	14.9	1.6	11.4	1.8	17.1	0.6	3.5	NS	NS
Number of deaths	175	370	105	394	70	816	1,930		

In DS, is the problem the lung, the immune system, or both?

The Lung is Overlooked in DS



The Lung is Overlooked in DS



Infectious lung disease accounts for 54% of hospital admissions for persons with DS

Average length of admission is 2-3 times longer than those without DS

Persons with DS have increased frequency of respiratory tract infection (62 fold higher rate)

Increased risk for acute respiratory distress syndrome

16 fold, 8 fold, 335 fold more likely to be hospitalized, intubated, or to die, respectively

Infectious respiratory disease accounts for more deaths in DS than any other medical condition; 12 times more likely to die than typical population

Lung Health is Front and Center for Persons with DS and Their Families

**Ever Made A Croup Tent At Home?
I Did Every Year From 1983-2004**



So What's the Mission, the Goal?

- 1. Awareness! Persons with DS, autoimmunity, & lung disease**
- 2. Learning how Trisomy 21 leads to respiratory & autoimmune diseases will help those with DS and those without DS**

Folks with DS will *feel better*-physically & mentally

Post-Influenza Pneumonia

Lessons from the Flu:

Changes in Immune Cell Function in Postinfluenza Bacterial Pneumonia

Flu usually not lethal, bacterial “super” infection often is (>50% of deaths)

Associated with high levels of interferons (types I and II), IL-10, TGF-beta

Type I IFNs mediate development of postinfluenza bacterial pneumonia in mice

Arash Shahangian,^{1,2} Edward K. Chow,³ Xiaoli Tian,⁴ Jason R. Kang,¹ Amir Ghaffari,^{1,2} Su Y. Liu,^{1,2} John A. Belperio,⁴ Genhong Cheng,^{1,5} and Jane C. Deng⁴

¹Department of Microbiology, Immunology and Molecular Genetics, and ²Medical Scientist Training Program, David Geffen School of Medicine, UCLA, Los Angeles, California, USA. ³G.W. Hooper Foundation, UCSF, San Francisco, California, USA.

⁴Division of Pulmonary and Critical Care Medicine and ⁵Molecular Biology Institute, David Geffen School of Medicine, UCLA, Los Angeles, California, USA.

Influenza-related complications continue to be a major cause of mortality worldwide. Due to unclear mechanisms, a substantial number of influenza-related deaths result from bacterial superinfections, particularly secondary pneumococcal pneumonia. Here, we report what we believe to be a novel mechanism by which influenza-induced type I IFNs sensitize hosts to secondary bacterial infections. Influenza-infected mice deficient for type I IFN- α/β receptor signaling (*Ifnar*^{-/-} mice) had improved survival and clearance of secondary *Streptococcus pneumoniae* infection from the lungs and blood, as compared with similarly infected wild-type animals. The less effective response in wild-type mice seemed to be attributable to impaired production of neutrophil chemoattractants KC (also known as Cxcl1) and Mip2 (also known as Cxcl2) following secondary challenge with *S. pneumoniae*. This resulted in inadequate neutrophil responses during the early phase of host defense against secondary bacterial infection. Indeed, influenza-infected wild-type mice cleared secondary pneumococcal pneumonia after pulmonary administration of exogenous KC and Mip2, whereas neutralization of Cxcr2, the common receptor for KC and Mip2, reversed the protective phenotype observed in *Ifnar*^{-/-} mice. These data may underscore the importance of the type I IFN inhibitory pathway on CXC chemokine production. Collectively, these findings highlight what we believe to be a novel mechanism by which the antiviral response to influenza sensitizes hosts to secondary bacterial pneumonia.

Observed in DS?



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YOU need not! Just carry Formamint with you and suck these delicious tablets whenever you are in danger of being infected by other people.

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“Attack the germs before they attack you!”

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Formamint
THE GERM KILLING THROAT TABLET

Important Mediator of the Enhanced Susceptibility to Bacterial Pneumonia after Influenza Infection

Sluijs,^{*,†‡} Leontine J. R. van Elden,[¶] Monique Nijhuis,[¶] Rob Schuurman,[¶] Adriene Florquin,[§] Michel Goldman,^{||} Henk M. Jansen,[†] René Lutter,^{†‡} and

pneumonia is a serious complication during and shortly after influenza infection. We established a mouse model for secondary pneumococcal pneumonia and evaluated the role of IL-10 in host defense against *Streptococcus pneumoniae* after influenza infection. C57BL/6 mice were intranasally inoculated with 10 median tissue culture ID₅₀ of influenza A (A/PR/8/34) or PBS (control) on day 0. By day 14 mice had regained their normal body weight and were free of viral infection, as determined by real-time quantitative PCR. On day 14 after viral infection, mice were challenged with *S. pneumoniae* (serotype 3) intranasally. Mice recovered from influenza infection were highly susceptible to secondary pneumococcal pneumonia, as reflected by a 100% lethality on day 3 after bacterial infection, whereas control mice showed only 3 and 83% lethality on day 6 after pneumococcal infection. Furthermore, 1000-fold higher bacterial loads were observed in mice with *S. pneumoniae* and, particularly, 50-fold higher pulmonary levels of IL-10 were observed in influenza-infected mice compared with control mice. Treatment with an anti-IL-10 mAb 1 h before bacterial inoculation resulted in a marked reduction in lethality during secondary bacterial pneumonia compared with those in IgG1-treated mice. These findings indicate that mild self-limiting influenza A infection renders normal immunocompetent mice highly susceptible to secondary pneumococcal pneumonia. This increased susceptibility to secondary bacterial pneumonia is at least in part caused by excessive IL-10 production and reduced neutrophil function in the lungs. *The Journal of Immunology*, 2004, 172: 7603–7609.

Hypothesis

In persons with DS, the lung is in a chronic state of susceptibility to severe *S.pneumoniae* pneumonia that phenocopies post-viral infection in non-DS individuals

Strong Inference



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SHORT REPORT



Interferonopathy

Toll-like receptors

Immune suppression IL-10/TGF-beta

Respiratory commensal bacteria

Trisomy 21 consistently activates the interferon response

Kelly D Sullivan^{1,2,3,4*}, Hannah C Lewis^{1,2}, Amanda A Hill^{1,2}, Ahwan Pandey^{1,2,3,4}, Leisa P Jackson^{1,3,4}, Joseph M Cabral^{1,3,4}, Keith P Smith¹, L Alexander Liggett^{1,5}, Eliana B Gomez^{1,3,4}, Matthew D Galbraith^{1,2,3,4}, James DeGregori^{1,5,6,7,8,9}, Joaquín M Espinosa^{1,2,3,4*}

J. theor. Biol. **86**, 603–606

LETTER TO THE EDITOR

Interferon Action and Chromosome 21 Trisomy

Hypothesis

In persons with DS, the lung is in a chronic state of susceptibility to severe *S.pneumoniae* pneumonia that phenocopies post-viral infection in non-DS individuals

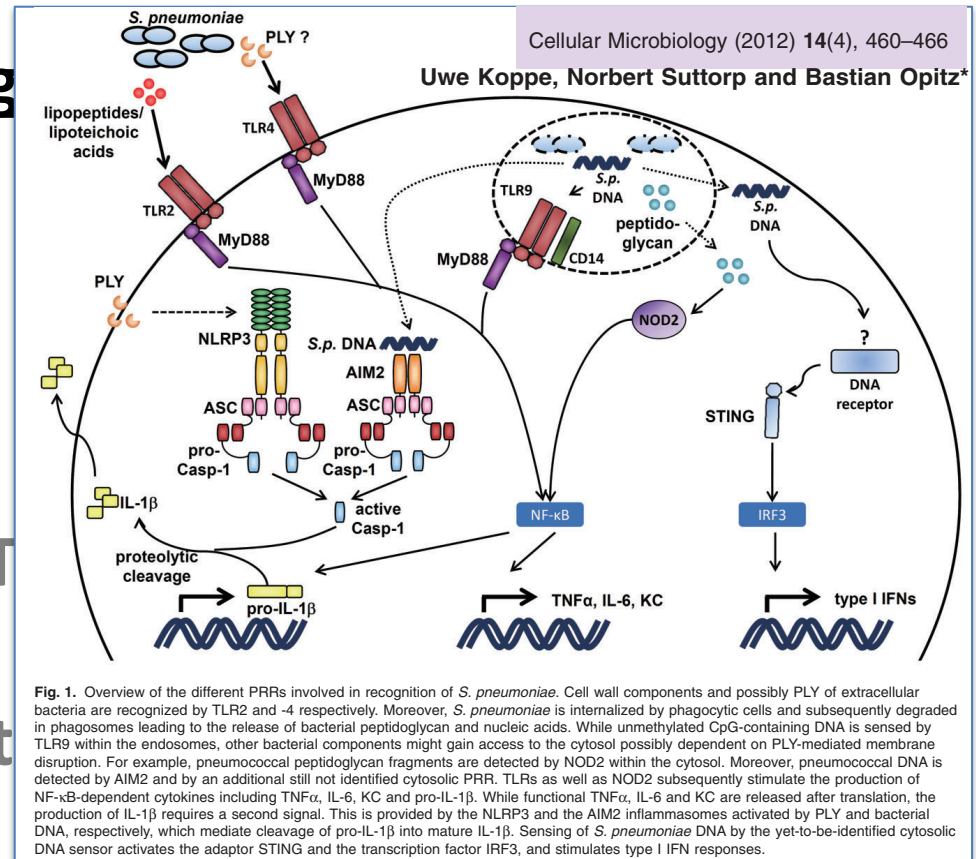
Strong

Interferonopathy

Toll-like receptors

Immune suppression IL-10/T

Respiratory commensal bact



Hypothesis

In persons with DS, the lung is in a chronic state of susceptibility to phenocopies p

Interferonopathy

Toll-like receptors

Immune suppressi

Respiratory comm

nature publishing group

Clinical Investigation

Articles

Increased production of interleukin-10 in children with Down syndrome upon *ex vivo* stimulation with *Streptococcus pneumoniae*

Chantal J.M. Broers¹, Reinoud J.B.J. Gemke¹, Servaas A. Morré^{2,3}, Michel E. Weijerman¹ and Anne Marceline van Furth⁴

BACKGROUND: Children with Down syndrome (DS) have an increased susceptibility to infections, due to altered humoral and/or cellular immunity. The aim of the study was to determine the cytokine production in whole blood of children with DS upon stimulation with heat-killed *Streptococcus pneumoniae* and lipopolysaccharide (LPS), in comparison with their healthy siblings.

METHODS: Whole blood of 61 children with DS and 57 of their healthy siblings was stimulated with 200 ng/ml LPS and 4×10^7 colony-forming units/ml *S. pneumoniae* during 6, 24, and 48 h. Concentrations of pro- and anti-inflammatory cytokines, tumor necrosis factor (TNF)- α , interleukin (IL)-1 β , IL-6, IL-8, IL-12p70, and IL-10 were determined at all time points.

RESULTS: Children with DS show an increased IL-10 production upon stimulation with *S. pneumoniae* compared to their healthy siblings. At most time points, no significant differences were seen in cytokine production upon stimulation with LPS.

CONCLUSION: Children with DS may be prone to a severe course of pneumococcal pneumonia, because of an increased anti-inflammatory response.

and the adaptive immunity are reported in DS, for example, mannan-binding lectin deficiency (13), a high number of pro-inflammatory CD14^{dim}CD16⁺ monocytes (14), changes in T- and B-lymphocyte counts (15–17), early aging of the immune system (18,19), an intrinsic defect of T and B lymphocytes (16,20,21), IgG2 and IgG4 subclass deficiencies (16,17,21–24), impaired antibody response to pneumococcal vaccine (25), and diminished invariant natural killer T cells (14,17) and regulatory T cells (17). These lower RTIs in DS children are most often caused by viral pathogens, such as respiratory syncytial virus. This can lead to severe respiratory syncytial virus bronchiolitis, a frequent cause of hospitalization in DS children (10,26–28). Also, an increased risk of hospitalization, endotracheal intubation, and death due to influenza A virus infection was reported in DS (29). In addition, we found an increased proinflammatory cytokine response to live influenza A virus in children with DS, which might contribute to an increased severity of their clinical course of this infection (30). Bacterial pathogens, both Gram positive and Gram negative, can also cause lower RTIs in children. However, nothing

Hypothesis

In persons with DS, the lung is in a chronic state of susceptibility to severe *S. pneumoniae* pneumonia that phenocopy



ORIGINAL RESEARCH
published: 23 August 2017
doi: 10.3389/fmicb.2017.01613



Respiratory Commensal Bacteria *Corynebacterium pseudodiphtheriticum* Improves Resistance of Infant Mice to Respiratory Syncytial Virus and *Streptococcus pneumoniae* Superinfection

OPEN ACCESS

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Julio Villena^{1,3,4*}

Interferonopa

Toll-like recep

Immune supp

Respiratory commensal bacteria is different

Post-Influenza-Like Lung in Down Syndrome?

Post-influenza, non-DS

DS

Fulminant Remodeling

?

Platelet Activating Factor Receptor (PAFR)

?

IFN signaling increase

IL-10 signaling increase

?

TLR down

?

Immune cell dysfunction

Post-Influenza-Like Lung in Down Syndrome?

Organ

Subpleural cysts

Pulmonary hypoplasia

Congenital heart defects

Tissue

Cells

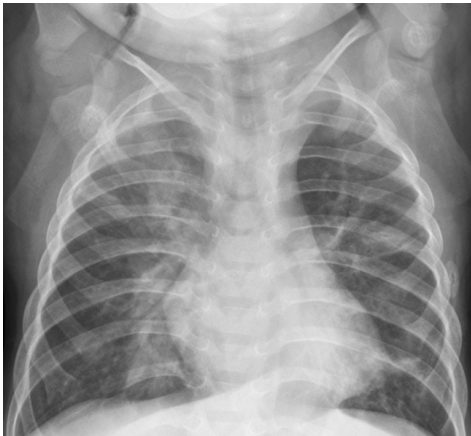


Fig. 5 Respiratory syncytial virus bronchiolitis. Anteroposterior radiograph of the chest in a 14-month-old girl with Down syndrome shows symmetrical hyperinflation of the lungs, streaky perihilar opacities and other focal right upper lobe opacities. The girl was diagnosed with respiratory syncytial virus (RSV) bronchiolitis



Fig. 9 Tracheal rings. Axial CT of the chest in a 23-month-old boy with Down syndrome shows a circular configuration of the trachea (*arrow*). This along with the small caliber of the trachea is consistent with a diagnosis of complete tracheal rings

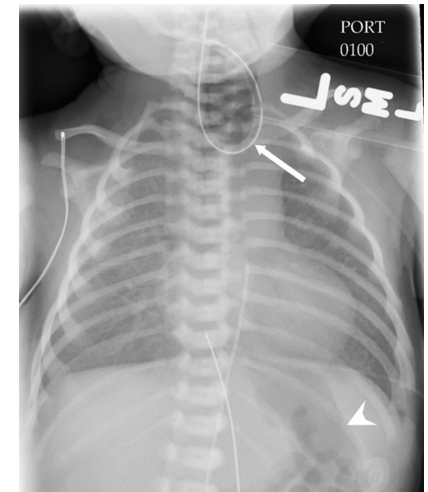


Fig. 3 Tetralogy of Fallot and esophageal atresia/tracheoesophageal fistula. Anteroposterior chest radiograph in a newborn boy shows that the heart has a boot-shaped contour with an upturned apex compatible with a diagnosis of tetralogy of Fallot. A nasogastric tube is coiled in the upper esophageal pouch (*arrow*), consistent with esophageal atresia. The presence of bowel gas (*arrowhead*) in the upper abdomen confirms the presence of a tracheoesophageal fistula. There are also 13 pairs of ribs

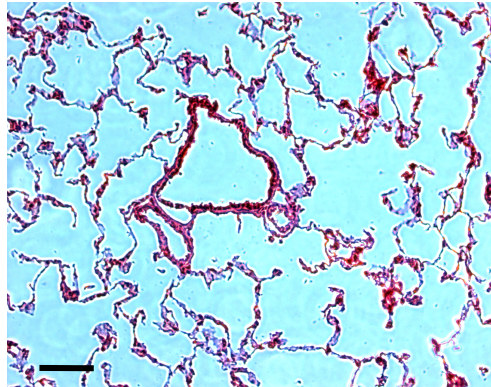
We need to explore mouse models of DS more fully for these

Post-Influenza-Like Lung in Down Syndrome?

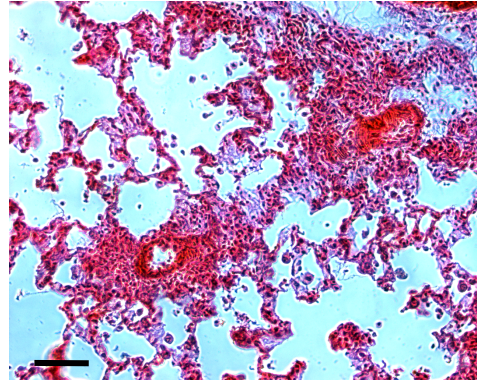
Organ

Tissue

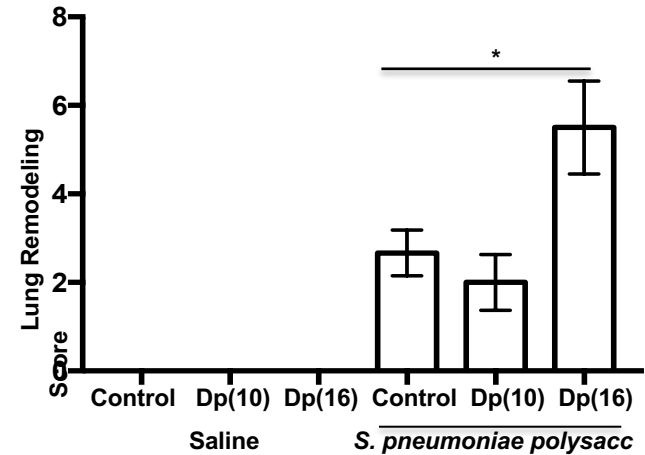
Cells



Saline



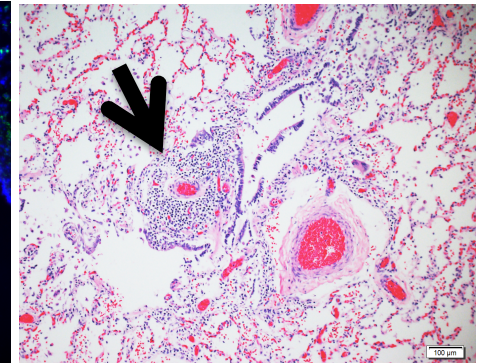
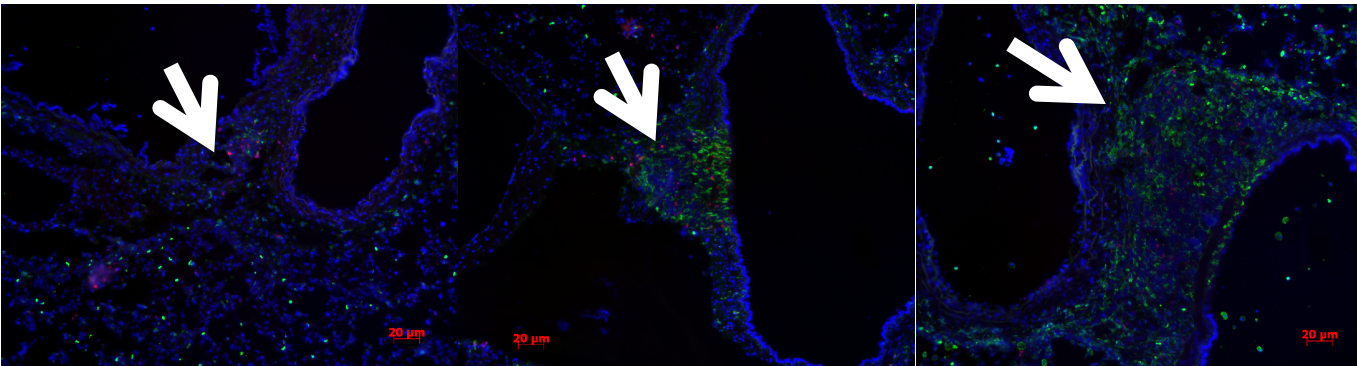
S. pneumoniae polysacc



B

Green = CD15 Blue = DAPI (nuclei)

Hematoxylin & Eosin



C

Control

Dp(10)

Dp(16)

DS

Post-Influenza-Like Lung in Down Syndrome?

Organ

Tissue

Increased PAFR facilitates increased *S. pneumo* adhesion

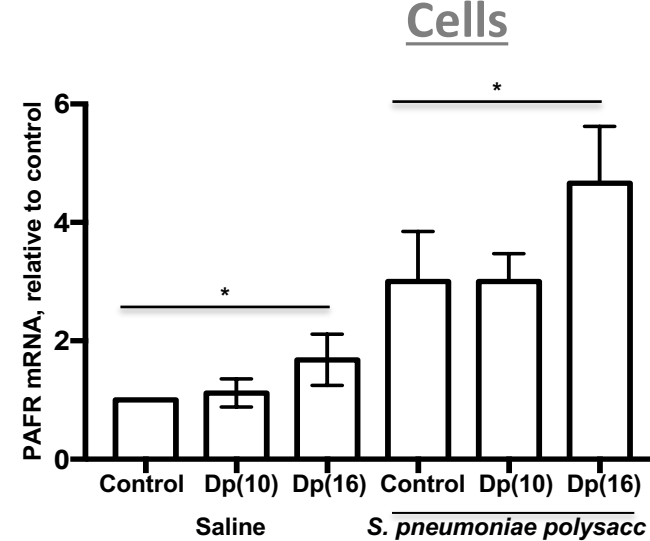
Involvement of the platelet-activating factor receptor in host defense against *Streptococcus pneumoniae* during postinfluenza pneumonia

Koenraad F. van der Sluijs,^{1,2,3} Leontine J. R. van Elden,⁴ Monique Nijhuis,⁴
Rob Schuurman,⁴ Sandrine Florquin,⁵ Takao Shimizu,⁶ Satoshi Ishii,⁷
Henk M. Jansen,² René Lutter,^{2,3} and Tom van der Poll¹

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⁶Department of Biochemistry and Molecular Biology, Faculty of Medicine, The University of Tokyo; and ⁷CREST of Japan Science and Technology Corporation, Tokyo, Japan

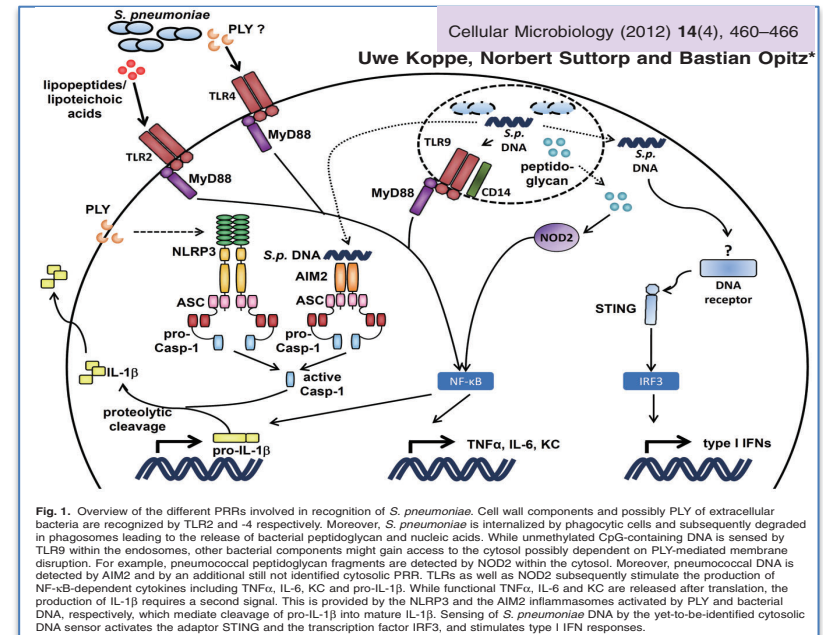
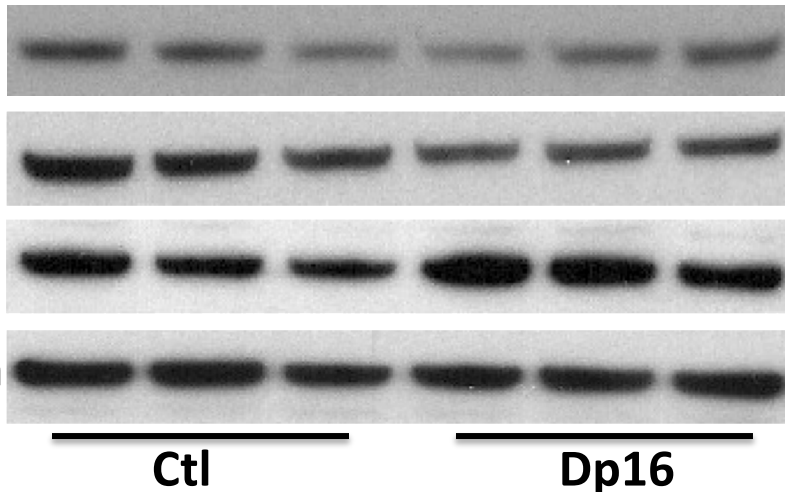


TLR2

TLR4

PAFR

Bactin



Post-Influenza-Like Lung in Down Syndrome?

Organ

Tissue

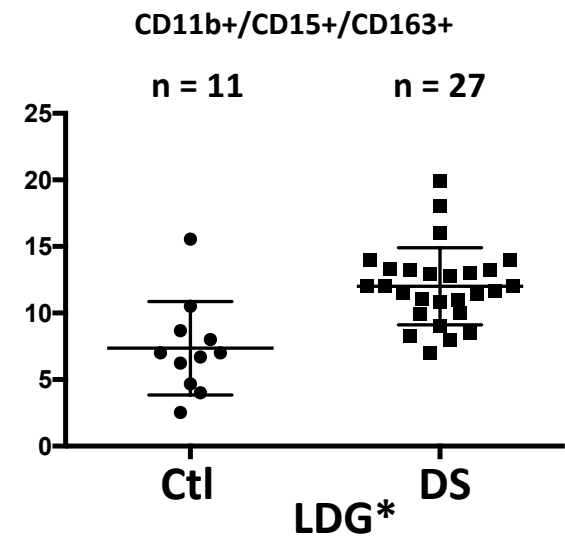
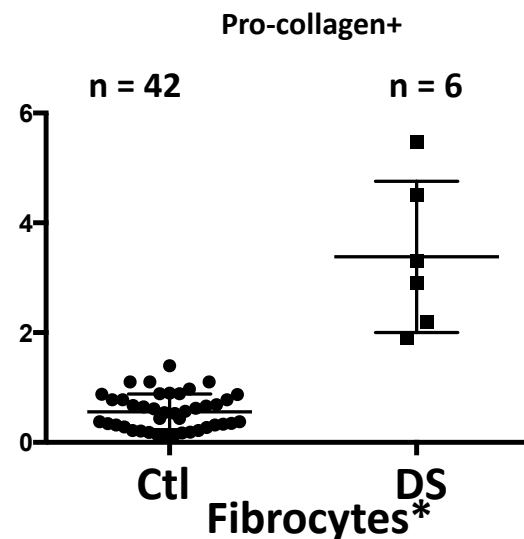
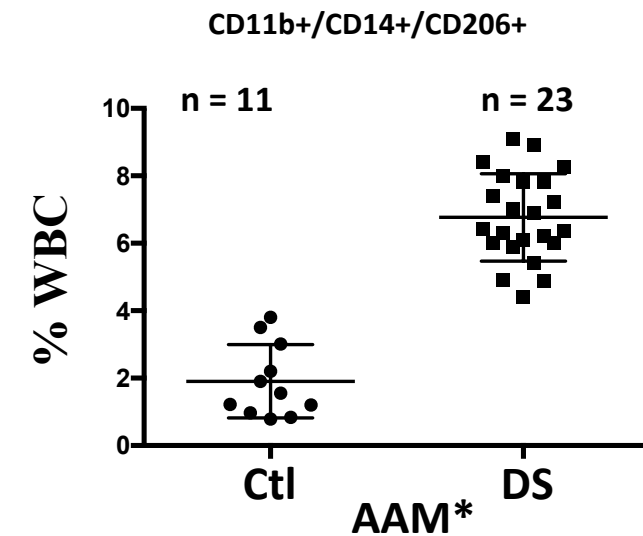
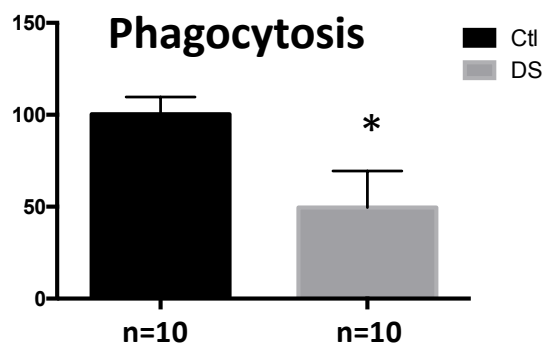
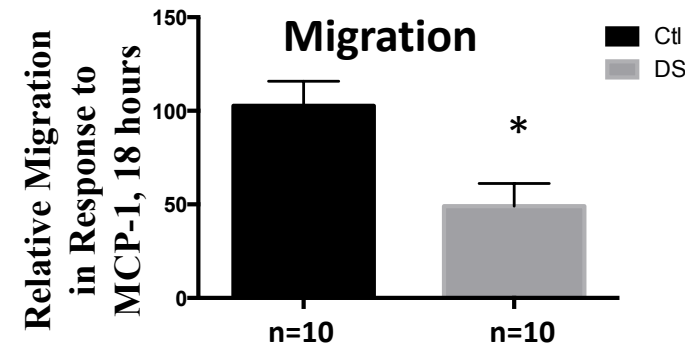
Cells

Lung stains

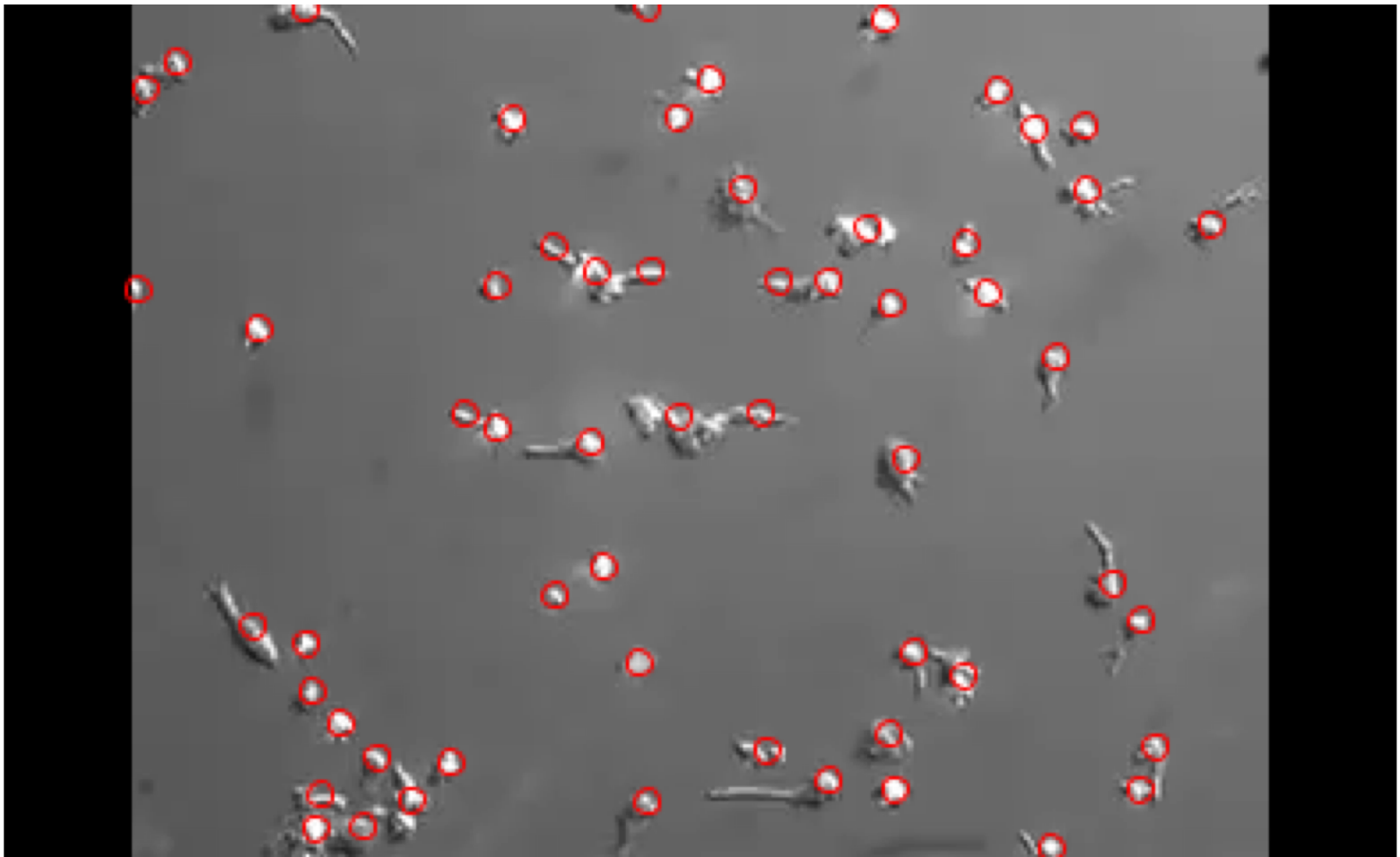
Human
Peripheral blood
Fibroblasts

Dp16
PB/BALF

Lung Cells



Cell/Individual particle tracking Icy



Post-Influenza-Like Lung in Down Syndrome?

Organ

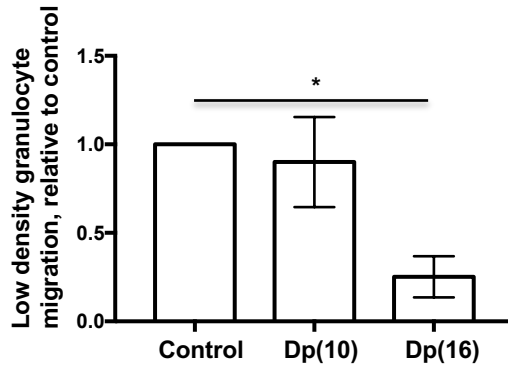
Tissue

Cells

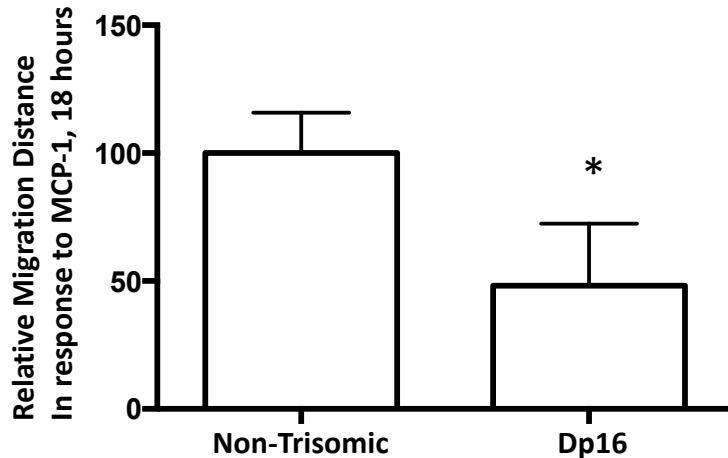
Lung stains

Human
Peripheral blood
Fibroblasts

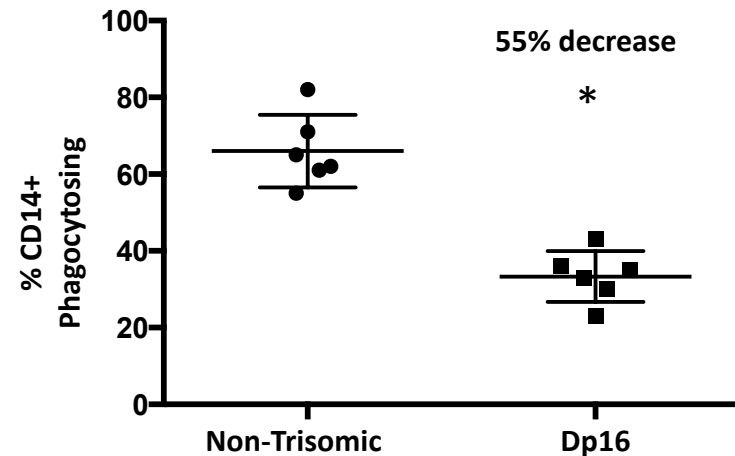
Dp16
PB/BALF
Lung Cells



Migration



Phagocytosis



Post-Influenza-Like Lung in Down Syndrome?

Post-influenza, non-DS

DS

Fulminant Remodeling

PAFR

IFN signaling increase

IL-10 signaling increase

TLR down

Immune cell dysfunction

Is there A Link Between Lung Disease and Cognition in Down Syndrome?

Child: care, health and development

Table 1. Patient characteristics and additional morbidity of 8-year-old Down syndrome population in relation to parent-reported presence of recurrent respiratory tract infections (RRTI)

Original Article

	RRTI ⁺	RRTI ⁻	Total	Chi-squared test
	n (%)	n (%)	n (%)	P-value
Male	85 (57)	84 (48)	169 (51)	NS
Female	64 (43)	92 (52)	156 (49)	
Age at inclusion* (mean, range and SD in years)	8.14 (7.8–8.8) ± 0.14	8.15 (7.8–9.1) ± 0.16	8.14 (7.8–9.1) ± 0.15	
School attendance				
Ever attended regular education	99 (30)	142 (44)	241 (74)	0.003
Regular education attendance at inclusion	65 (21)†	91 (28)	156 (48)	NS
Pre-school (normally age 1–5 years)	21 (32)†	11 (12)	31 (20)	0.018
First grade (normally age 6 years)	33 (51)†	62 (67)	95 (61)	0.010
Second grade (normally age 7 years)	11 (17)†	17 (19)	28 (19)	NS
Level of parental education				
Primary or secondary education	24 (7)	31 (10)	55 (17)	NS
Higher secondary education	55 (17)	63 (19)	118 (36)	NS
University education	70 (22)	82 (25)	152 (47)	NS
Being fostered (1 month)	57 (18)	60 (18)	117 (36)	NS
Siblings	140 (43)	170 (52)	310 (95)	NS
Child was born at home	142 (44)	160 (49)	302 (93)	NS
Additional morbidity‡				
Congenital heart disease	73 (49)	64 (36)	137 (42)	0.022
Diagnosis of asthma	28 (19)	6 (3)	34 (10)	<0.001
Gastrointestinal disease	20 (17)	19 (11)	39 (14)	NS
Eye disease	77 (52)	81 (46)	158 (49)	NS
Impaired hearing	66 (44)	32 (18)	98 (30)	<0.001
Thyroid dysfunction	19 (13)	20 (11)	39 (12)	NS
Diabetes mellitus	1 (<1)	2 (1)	3 (1)	NS
Other morbidity, not specified	43 (29)	38 (22)	81 (25)	NS

*There was no significant difference in age between both groups determined by a *t*-test.

†Percentage out of all children attending regular education.

‡Parental reported morbidity.

RRTI⁺, children with respiratory tract infections; RRTI⁻, children without respiratory tract infections; NS, not significant.

Significant impact of recurrent respiratory tract infections in children with Down syndrome

R. H. J. Versteegen,* H. B. M. van Gameren-Oosterom,† M. Fekkes,† E. Dusseldorp,† E. de Vries* and J. P. van Wouwe†

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Accepted for publication 24 April 2012

Is there A Link Between Lung Disease and Cognition in Down Syndrome?

Table 2. Results of multiple regression analyses for scale scores of the McCarthy Scales of Children's Abilities (MSCA) of 8-year-old Down syndrome children with and without parent-reported recurrent respiratory tract infections (RRTI)

	RRTI ⁺		RRTI ⁻		Regression coefficient [†] (β)	Effect size [‡] (f^2)
	Total (n = 130)	Male (n = 75) Female (n = 55)	Total (n = 140)	Male (n = 69) Female (n = 71)		
Verbal	33.06 (19.13) [§]	29.64 (19.96) 37.47 (17.18)	40.68 (16.18)	37.09 (15.72) 44.11 (16.08)	-7.33**	0.04
Perceptual performance	26.76 (16.58)	21.80 (15.84) 33.33 (15.43)	32.34 (14.90)	28.83 (14.87) 35.93 (14.20)	-5.67**	0.03
Quantitative	9.44 (6.80)	7.86 (6.46) 11.44 (6.75)	12.22 (6.57)	10.46 (6.75) 13.93 (5.99)	-2.61**	0.04
Memory	10.74 (7.77)	9.27 (7.69) 12.64 (7.53)	13.94 (7.37)	11.77 (6.31) 16.10 (7.77)	-3.12**	0.04
Motor	23.23 (12.57)	20.03 (12.64) 27.47 (11.32)	27.67 (11.78)	24.99 (11.51) 30.25 (11.62)	-4.63**	0.04
General cognitive score	69.30 (40.25)	59.36 (40.22) 82.29 (36.84)	85.24 (34.43)	76.35 (33.60)	-15.59**	0.04
Developmental age (SD in months)	3 years 8 months (10.91)	3 years 6 months (10.53) 3 years 11 months (10.66)	4 years 0 months (10.91)	4 years 0 months (10.91)		

Lower scores indicate more impaired development.

* $P < 0.05$; ** $P < 0.01$; *** $P < 0.001$.

[†] β = unstandardized regression coefficient of the effect of RRTI, correcting for age (>1 month), siblings, gender, congenital heart defect, diagnosis of asthma, gastroesophageal reflux.

[‡]Effect size (f^2): small effect (0.01–0.10), moderate effect (0.10–0.33) and large effect (>0.33).

[§]Mean scores are presented with standard deviation between brackets.

RRTI⁺, children with recurrent respiratory tract infections; RRTI⁻, children without recurrent respiratory tract infections.

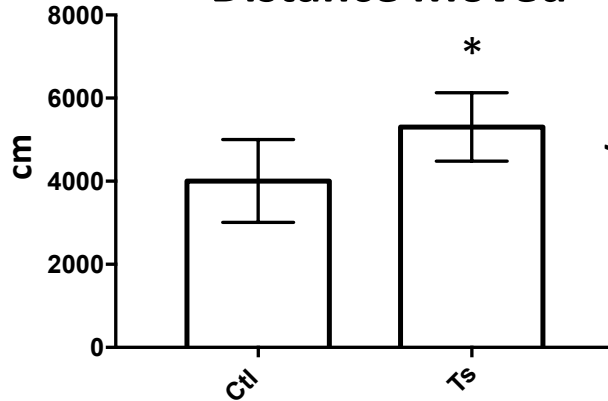
Key messages

- Children with Down syndrome are known to be at increased risk of recurrent respiratory tract infections.
- In 8-year-old children with Down syndrome, parental report of recurrent respiratory infections was associated with more delayed development, increased risk of behavioural problems and lower health-related quality of life.

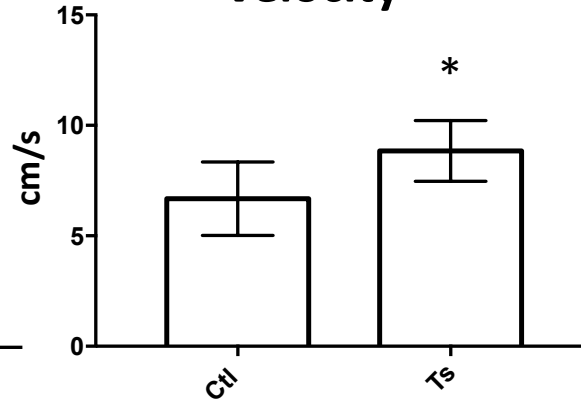
Lung Disease & Cognition: Tantalizing Clues

Cognitive Assays in Dp16 mice

Distance Moved

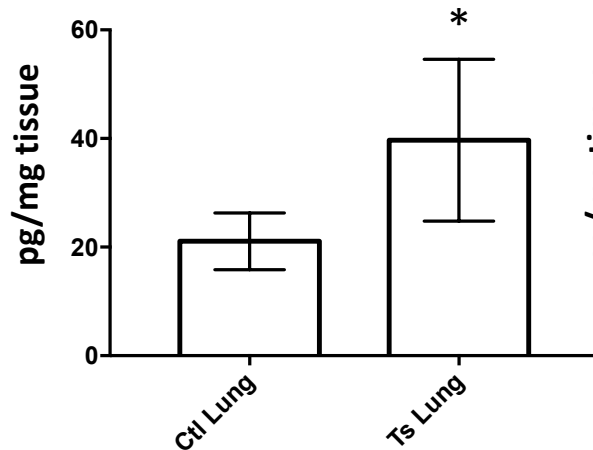


Velocity

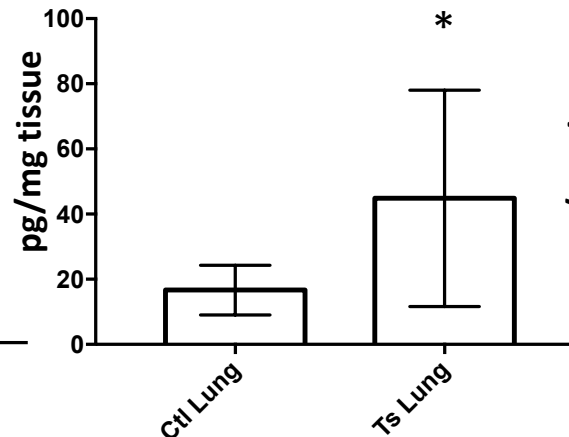


Do lower learning and memory scores correlate to inflammatory cytokines?

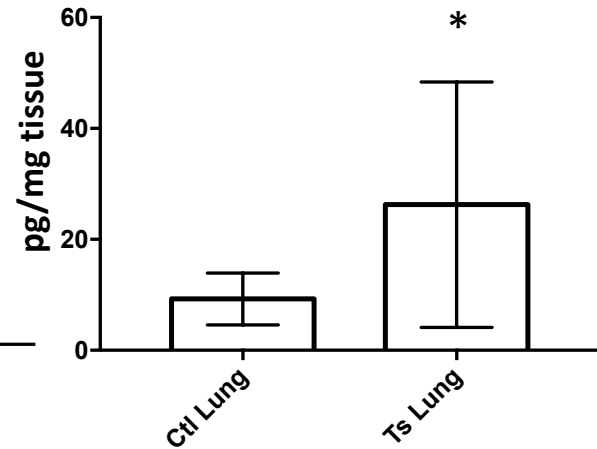
TNFalpha



IL-1b



Do repeat lung infections worsen cognitive score?



Take Home Messages

Individuals with DS are significantly challenged by infectious lung disease

Post-influenza state of susceptibility to severe bacterial lung infection

Chronic respiratory infection linked to myositis/myopathy, myocarditis, CNS inflammation

Infectious lung disease likely impacts cognition

Reducing burden in DS would greatly improve QOL

Reducing burden in DS may preserve cognition

The future is BRIGHT!! Great things have happened, and MUCH more is on the way

ex: Amniotic fluid stem cells “trained” to patch congenital heart defects-submit July 23rd

ex: Autoimmunity and Lung Disease

Many Thanks

Kelley Colvin

Persons with DS and
their families

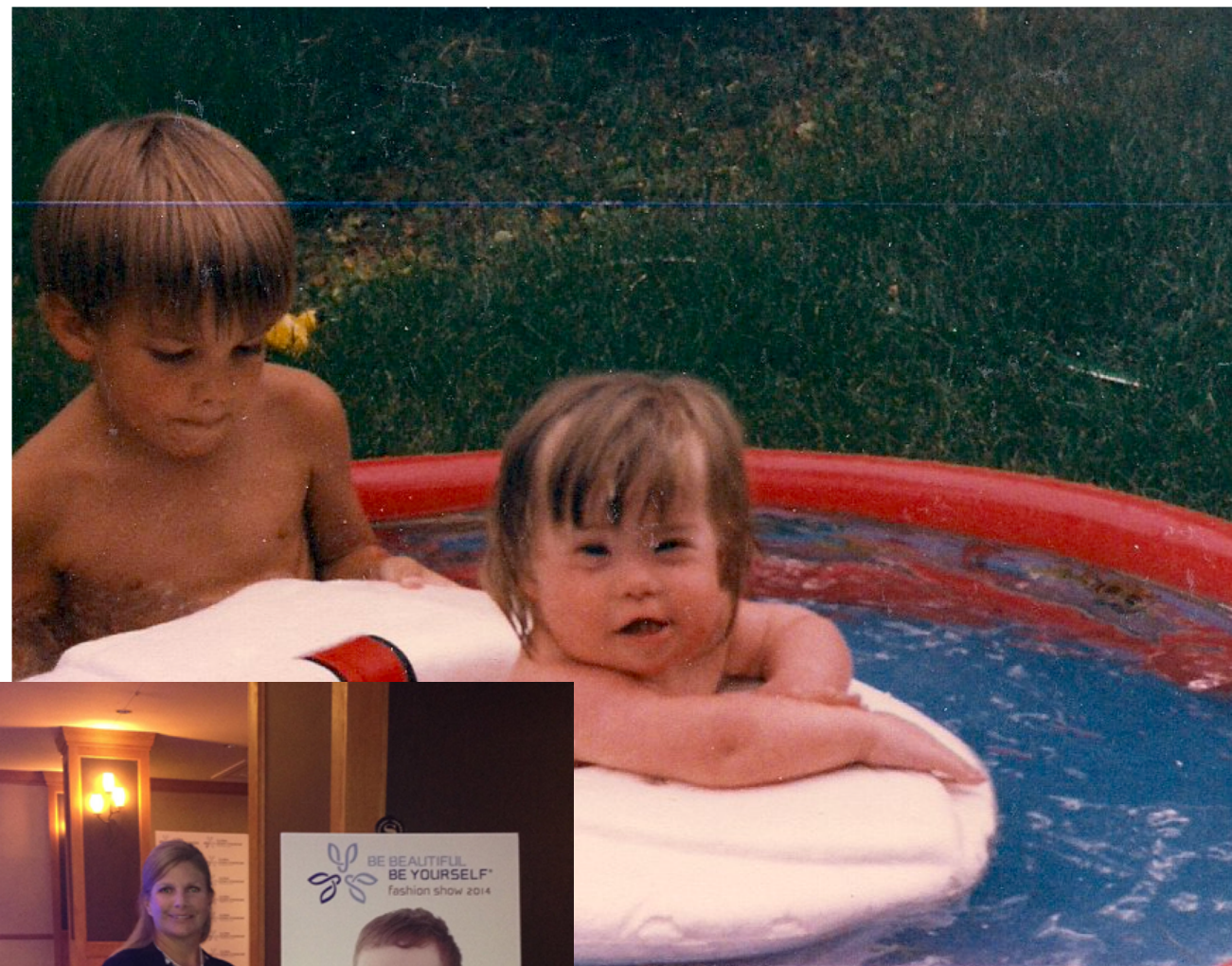
Support

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